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Mesoporous silica formation mechanisms probed using combined Spin-Echo Modulated Small Angle Neutron Scattering (SEMSANS) and Small Angle Neutron Scattering (SANS)

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Abstract

The initial formation stages of surfactant-templated silica thin films which grow at the air-water interface were studied using combined Spin-Echo Modulated Small-Angle Neutron Scattering (SEMSANS) and Small Angle Neutron Scattering (SANS). The films are formed from either a cationic surfactant (CTAB) or non-ionic surfactant (C₁₆EO₈) in a dilute acidic solution, by addition of TMOS. Previous work has suggested a two stage formation mechanism with mesostructured particle formation in the bulk solution driving film formation at the solution surface. From the SEMSANS data, it is possible to pinpoint accurately the time associated with the formation of large particles in solution, that go on to form the film, and to show their emergence is concomitant with the appearance of Bragg peaks in the SANS

pattern, associated with the 2D-hexagonal order. The combination of SANS and SEMSANS allows a complete depiction of the steps of the synthesis that occur in the subphase.

Keywords: Small angle neutron scattering (SANS), spin-echo modulated small angle neutron scattering (SEMSANS), mesoporous silica, kinetics, micelles, mesophase, particles, silica films

Introduction

Mesoporous materials, discovered in 1992,¹ present a well-defined and organised porosity. They are formed by mixing an amphiphilic structuring agent with an inorganic precursor in solution. Depending on the synthesis conditions, the material is synthesized as sub-micrometric grains in solution,²⁻³ monoliths,⁴⁻⁵ or as thin-films, either via spin-coating,⁶ dip-coating⁷⁻⁸ or by spontaneous formation at the air-water interface.⁹

Ever since the groups of Ozin¹⁰ and Aksay¹¹ reported on the synthesis of mesophase inorganic thin films formed at the solution surface in 1996, these materials have been a topic of great interest. The ability to manipulate the mesostructure in the films, their intrinsic porosity, the size of their pores, and other morphological parameters give the films potential applications in a wide variety of scientific and technological fields such as selective membranes,¹² catalysis,¹³⁻¹⁴ gas sensors,¹⁵ and drug-delivery.¹⁶ Extensive research has been done on these self-assembling silica-surfactant systems; however, the mechanism of this self-assembly, leading to the formation of the mesostructured thin films, is still a matter of debate. Currently, the two possible mechanisms that have been proposed in the literature are the silica-coated micelles mechanism and the silica-droplet phase separation mechanism.⁹ The former relies on the addition of silica species to the corona of the micelles in the solution, influencing the micelles shape, usually promoting more elongated packing parameters. Then, either some of the silica-enriched micelles migrate to the interface to form the film, or alternatively form

mesostructured particles in the subphase that later migrate to the interface. The latter mechanism suggests the rapid formation of phase-separated droplets rich in surfactant and silica species - often called flocs - and possibly remaining-quasi globular micelles in the bulk. The flocs are suggested as nucleation sites for micelle elongation and reorganisation into a mesostructure. In the first case, the signal of elongating micelles should be measured in the bulk before the signal of larger flocs, while the second hypothesis suggests the appearance of large objects in the bulk prior to any reshaping of the micelles.

To investigate the behaviour of the micelles and particles in the bulk, a combination of traditional Small Angle Neutron Scattering (SANS) and Spin-Echo Modulated Small-Angle Neutron-Scattering (SEMSANS)¹⁷⁻¹⁹ was implemented on the Larmor instrument (ISIS, Rutherford Appleton Laboratory, UK). This allowed for the investigation of phenomena on the nano- and micro-metre scale simultaneously, providing valuable insights on the behaviour of the micelles prior to and during aggregation. Furthermore, it enables the identification of the formation of an ordered mesostructure within the larger aggregates. Finally, when evidence was found for the phase-separation mechanism, SANS/SEMSANS was used to determine whether the micelles aggregated in an orderly fashion, or whether random aggregation was followed by ordering into the later observed mesostructure. Because this was the first-ever application of this combined technique on a real physical-chemical system, a description of the setup is provided below.

Material and Methods

Films preparation

Mesoporous silica film forming solutions were prepared following already established protocols.²⁰⁻²¹ The films are formed from solution by mixing tetramethoxysilane (TMOS) with a surfactant solution in acidic conditions. The surfactant (SF) used was either

hexadecyltrimethylammonium bromide (CTAB) or octaethylene glycol mono-hexadecyl ether (C₁₆EO₈) depending on the synthesis. The C₁₆EO₈ was purchased from Nikko Chemicals, Japan (BC-8SY). The CTAB, HCl, D₂O and TMOS were purchased from Sigma-Aldrich and all were used as received. For each synthesis, the molar ratio SF:HCl:D₂O was 1:2.39:658, in agreement with previous studies.⁹ Several syntheses were studied by varying the molar ratio between the silica source and the surfactant. In detail, three molar ratios $r_{mol} = \frac{n_{TMOS}}{n_{SF}} = 3.6$; 7.21 and 10.81 were investigated, using CTAB as surfactant, and one ratio ($r_{mol}=7.21$) for C₁₆EO₈. Those ratios correspond to ratios previously investigated with other techniques.⁹ TMOS was added to the already prepared surfactant solution in 0.2M HCl, which was then stirred for one minute before filling the SANS cell. The cell cap was left off to allow evaporation and hence have similar synthesis conditions to previous studies.

Combined SEMSANS and SANS measurements

In a typical small angle neutron scattering (SANS) measurement, the sample loaded in a cell is exposed to a well-collimated neutron beam, inducing scattering. The intensity of the scattered beam I is recorded on a pixelated detector, while the non-scattered, direct, beam is discarded using a beamstop. The scattered intensity is azimuthally averaged and given as a function of the scattering vector $q = \frac{4\pi}{\lambda} \sin(\frac{\theta}{2})$, with λ the wavelength of the neutron beam and θ the scattering angle. The q -range accessible in SANS permits probing dimensions typically ranging from 1 to a few hundred nm.

To access the larger μm -scale structures at very low q , we have to measure the scattering which is contained within the footprint of the direct beam, which is usually discarded. However, with the addition of several components we can gain additional information using a new technique called spin-echo modulated small angle neutron scattering (SEMSANS), proposed by Gähler.²² We implemented this method using a polarised beam and two

superconducting Wollaston prisms²³⁻²⁴ to create a spatial modulation of the intensity of the neutron beam at the detector position. The change in modulation amplitude caused by scattering from the sample gives the sample density autocorrelation function $G(\delta_{SE})$,¹⁷⁻¹⁹ where δ_{SE} , the spin-echo length, is the length scale being probed and is dependent upon the period (p) of the neutron intensity modulation at the detector. The quantity $G(\delta_{SE})$ is the same as that measured in spin-echo small angle neutron scattering (SESANS)²⁵ and we can use the same models and transforms to interpret the data. Unlike SESANS, the SEMSANS method has all the components for neutron spin manipulation before the sample and hence can be combined with the SANS technique for simultaneous measurements. Due to the use of time of flight, a range of wavelengths is probed at the same time, allowing measuring SEMSANS and SANS simultaneously.

A detailed description of the different components of the setup for the SEMSANS measurements, and of the procedure for the data corrections can be found in refs²⁶⁻³¹ and in the supplementary information.

The measurements were performed on the LARMOR instrument at the ISIS Pulsed Neutron and Muon Source. The sample to detector distance is 4 m and the accessible SANS q -range is 0.08-0.4 Å⁻¹, while the accessible range of spin-echo lengths was 0.35 < δ_{SE} ≤ 2.5 μm. A series of runs were performed every 19 minutes until an equilibrium state was found.

SESANS

In addition to the combined SEMSANS/SANS measurements, we also utilised the monochromatic SESANS instrument at the TU-Delft.³² Briefly, this utilises a 2.03 Å beam with a series of inclined magnetised films to encode the polarisation. This instrument has encoding films before and after the sample and cannot be combined with SANS. However, this is able to reach significantly higher spin-echo lengths. For these measurements the same

correlation function $G(\delta_{SE})$ is determined. Nonetheless, measurement at each spin-echo length has to be performed separately (due to the monochromatic nature of the beam) and the sample was measured for eleven different spin-echo lengths which were cycled through every 19 minutes (i.e. 100 s measurement per spin-echo length). This was to preserve a similar time structure to the combined SEMSANS/SANS measurements made on LARMOR.

SANS data treatment

The theoretical model for the formation of mesostructured materials studied in small angle scattering has already been described in the literature.³³⁻³⁴ The expression for the intensity can be described as the sum of four terms:

$$I_{tot}(q) = I_{micelles}(q) + I_{Bragg}(q) + I_{Porod}(q) + bg \quad (1)$$

with $I_{micelles}(q)$ the contribution of the ellipsoidal micelles of surfactant in solution; $I_{Bragg}(q)$ the contribution of the forming mesostructured material; $I_{Porod}(q)$ a Porod term at the very low angles associated with the forming particles and bg a constant background term.

For ellipsoidal micelles with radii $(R, R, \varepsilon R)$, $I_{micelles}(q)$ (in cm^{-1}) can be expressed as:³⁵

$$I_{micelles}(q) = AP(q, R, \varepsilon, \sigma) \quad (2)$$

with $A = n(\rho_{micelles} - \rho_0)^2$ the scale factor, depending on the number of micelles per unit volume n (in cm^{-3}), $\rho_{micelles}$ and ρ_0 the scattering length densities of the core of the micelles and solvent respectively ($\rho_0 = 6.335 \times 10^{10} \text{ cm}^{-2}$ for D_2O), and $P(q)$ the form factor of the micelles with their volume included (in cm^6). $P(q)$ is defined as:

$$P(q, R, \varepsilon, \sigma) = \langle \int_0^{\pi/2} [F(q, r(R, \varepsilon, \theta))]^2 \sin\theta d\theta \rangle_\sigma \quad (3)$$

with $r(R, \varepsilon, \theta) = R(\sin^2\theta + \varepsilon^2 \cos^2\theta)^{1/2}$, to describe the ellipticity of the radii.

$$F(q, r) = V(r)f_s(q, r) \quad (4)$$

is the form factor amplitude for a sphere of core radius r . $V(r) = \frac{4}{3}\pi r^3$ is the volume of a sphere of radius r and $f_s(q, r) = \frac{3[\sin(qr) - qr\cos(qr)]}{(qr)^3}$ the normalized form factor amplitude of a sphere of radius r .

Finally, the polydispersity in size $\langle . \rangle_\sigma$ is taken into account using the Schulz-Zimm distribution, i.e, for a certain function $F'(q, r)$:

$$\langle F'(q, r) \rangle_\sigma = \int f_{SZ}\left(R, r, \frac{1}{\sigma^2} - 1\right) F'(q, R) dR \quad (5)$$

$$\text{with } f_{SZ}(x, x_0, z) = \frac{x^z}{\Gamma(z+1)} \left(\frac{z+1}{x_0}\right)^{z+1} e^{-(z+1)\frac{x}{x_0}}$$

In order to easily compare the contribution from the free micelles to the overall intensity, one can rewrite equation 2 as follow:

$$I_{micelles}(q) = I_{micelles}(q=0)P'(q, R, \varepsilon, \sigma) \quad (6)$$

with

$$\begin{aligned} I_{micelles}(q=0) &= AP(0, R, \varepsilon, \sigma) = n(\rho_{micelles} - \rho_0)^2 P(0, R, \varepsilon, \sigma) \\ &= n(\rho_{micelles} - \rho_0)^2 V_{ell}^2(R, \varepsilon) \end{aligned} \quad (7)$$

the intensity from the micelles at $q=0$ (in cm^{-1}) and $P'(q, R, \varepsilon, \sigma) = \frac{P(q, R, \varepsilon, \sigma)}{P(0, R, \varepsilon, \sigma)}$ the normalised form factor. $V_{ell}(R, \varepsilon) = \frac{4}{3}\pi \varepsilon R^3$ is the volume in cm^3 of an ellipsoid of radius $(R, R, \varepsilon R)$.

We note that previous studies, using different $\text{D}_2\text{O}/\text{H}_2\text{O}$ contrasts, showed that the micelles are of a core-shell type.³⁶ Nonetheless, in our case, with only one contrast, we can fit the scattering from the micelles using only this homogeneous ellipsoid form factor.

The contribution associated with the mesophase is given by $I_{Bragg}(q)$:³⁷⁻³⁸

$$I_{Bragg}(q) = \sum_{hkl} I_{hkl} L_{hkl}(q, a, D, \nu) \quad (8)$$

with $L_{hkl}(q, a, D, \nu)$ the function that will reproduce the lattice organisation (e.g. Bragg peaks found at the positions q_{hkl}), taking into account the lattice parameter a (in nm) associated to the peak position and type of organisation, the lattice dimension D (in nm) which depends on the peak width, the numerical parameter ν associated with the peak shape (0 for a Lorentzian shape, 1 for a Gaussian shape), and l, h, k the Miller indices of the reticular planes associated to the mesophase. Values of the h, k, l indices, and hence the peaks positions will be highly dependent on the type of mesophase studied. In the q -range used in this study, only the first peak can be measured, which prevents discrimination of which type of mesophase is forming (2D hexagonal, cubic etc) and hence extraction of the lattice parameter a .

Nonetheless, for the CTAB templated films, we can assume that the Bragg peak for the particles formed in the subphase is associated with a 2D-hexagonal ordering, as already well-established in the literature^{36, 39-40} and measuring one peak is enough to extract both the lattice parameter and lattice dimension. Indeed, for a 2D-hexagonal mesophase, the position of the Bragg peaks is given by the formula $q_{hkl} = q_{hk} = \frac{4\pi}{a\sqrt{3}}\sqrt{h^2 + k^2 + hk}$. The first peak is in that case associated with the (10) plane and is found at position $q_{10} = \frac{4\pi}{a\sqrt{3}}$. The intensity parameter of this peak I_{10} can be written:

$$I_{10} = I(q = q_{10}) = n'(\rho_{micelles} - \rho'_0)^2 P(q_{10}, R, \varepsilon, \sigma) \quad (9)$$

It is associated not only with the number of micelles within the mesostructured phase domains n' , but also the contrast between the micelles and the silica walls $(\rho_{micelles} - \rho'_0)^2$, where ρ'_0 is the scattering length density of the condensing “silica walls” between the micelles). As

only one Bragg peak was extracted, it is not possible to separate the two parameters n' and ρ'_0 .

The films templated with C₁₆EO₈ form a variety of mesophases dependant on the silica:surfactant molar ratio. At the ratio studied here, $r_{mol}=7.2$, the films contain a $Pm\bar{3}n$ cubic phase,²¹ however the structure within the particles in the subphase cannot be determined from the single peak observed here, although, as for the CTAB templated particles, it can still be used to extract a d-spacing and domain size.

Finally we note that the width of the peak is also limited by the resolution in q (called Δq) of the SANS detector. This means the domain size of the mesophase can be accurately extracted only for values $D \leq \frac{2\pi}{\Delta q}$.

At small angles, the Porod term $I_{Porod}(q)$ is associated with the growing mesophase, with a parameter related to the smooth interface of the forming mesostructured particles:

$$I_{Porod}(q) = \frac{K}{q^4} \quad (10)$$

The q -range used in SANS does not allow extraction of the size of the particles, which is instead observed using the spin-echo configuration.

SEMSANS and SESANS data treatment

For both SEMSANS and SESANS, data are expressed in the form of $(G(\delta_{SE}) - 1) * \Sigma_t = \frac{\ln(P/P_0)}{\lambda^2 t}$ as a function of spin-echo length (δ_{SE}). where, λ is the neutron wavelength, t the sample thickness and Σ_t a measure of the scattering power. In SEMSANS, $P=P(\lambda)$ is the amplitude of the intensity modulations on the detector for the sample at a given wavelength, while P_0 is the one from the blank; while in SESANS they are the amplitude of the

polarisation directly. A more detailed explanation of the extraction of $\frac{\ln(P/P_0)}{\lambda^2 t}$ versus δ_{SE} is given in S.I and in ref ⁴¹.

Both the SESANS and SEMSANS data were fitted to a simple dilute sphere model as given by Andersson et al.²⁵ This is characterised by two parameters: a measure of the scattering power independent from wavelength and sample thickness (Σ_t) and radius ($R_{particles}$ in μm). We note that, for concentration of particles below ca 10%:

$$\Sigma_t = \frac{3}{2} R_{particles} (\Delta\rho)^2 \varphi (1 - \varphi) \quad (12)$$

with $\Delta\rho$ the contrast between the particles and the solvent and φ the volume fraction of particles. This means that Σ_t and $R_{particles}$ are correlated. In the plots of $\ln(P/P_0)/(\lambda^2 t)$ vs δ_{SE} (see for example Figure 1.b), the signal decreases until it reaches a plateau. The position in δ_{SE} of the plateau can be related to $2R_{particles}$, while the value of $\ln(P/P_0)/(\lambda^2 t)$ at the plateau is related to Σ_t . In SEMSANS, we can only probe a limited spatial range of spin-echo lengths δ_{SE} ($\leq 2.5 \mu\text{m}$), which means we will have an accurate measurement of $R_{particles}$ and Σ_t only for particles $\leq 2.5 \mu\text{m}$, where the plateau of the signal is clearly observed. This is nonetheless sufficient to evidence the formation of μm -sized particles in this study.

Results

Synthesis using CTAB at a ratio $r_{mol}=3.6$

A synthesis solution made using CTAB as surfactant with a molar ratio $r_{mol}=3.6$ was investigated using the combined SEMSANS and SANS setup, with SANS and SEMSANS patterns as a function of time given in Figure 1. This ratio has the lowest amount of silica of the three investigated, hence presents a slower speed of reaction. Both the SANS and the SEMSANS data were fitted using the models for mesostructured 2D-hexagonal domain

growth kinetics and for growth of spherical particles respectively, detailed in the Materials and Methods section. SANS patterns (Figure 1.a) demonstrate that at the beginning of the reaction, CTAB micelles in solution are present, with a slightly prolate shape. Over time, an elongation of the micelles is observed, before a Bragg peak arises at around 2 hours (observed but too weak to be fitted at 110 min, labelled below as t_{Bragg}), associated with the formation of a hybrid 2D-hexagonal mesostructure, which further grows in intensity with time. Concomitantly to the Bragg peak's initial appearance, the SEMSANS signal diverges from a flat line (Figure 1.b, divergence is present at 110 min but clearer at 129 min), with a modulation observed around $\delta_{SE}=1.5\text{ }\mu\text{m}$, characteristic of particles typically bigger than $1\text{ }\mu\text{m}$ in solution.

A detailed analysis of the parameters extracted from fitting the SANS and SEMSANS data for the entire reaction is given in Figures 2 and 3.

Figure 2 allows a clear comparison with time between $I_{micelles}(q=0)$ the intensity from the micelles of CTAB in solution, I_{10} the peak intensity associated with the mesophase organisation and Σ_t the scattering power associated with the μm -sized particles observed in SEMSANS. The signal associated with the micelles firstly increases until $t_{Bragg}=110\text{ min}$, before strongly decreasing while signals for both mesostructure and μm -sized particles increase after t_{Bragg} . The first increase of $I_{micelles}(q=0)$ is assumed to be due to the incorporation of silica species into the CTAB micelles, modifying both the micelle shape and their contrast with the solvent as further described in the next paragraph. The decrease after the emergence of the Bragg peak is explained by the continuing consumption of the micelles to form the mesophase. On the other hand, signals associated with the 2D-hexagonal phase and the μm -sized particles appear around the same time (although I_{10} was too weak to be fitted at 110 min). Σ_t shows a sharp increase before fluctuating around a plateau, which seems to indicate that large particles are quickly formed and then their number remains roughly

constant with time. Comparatively, the signal of I_{10} grows more slowly upon time, mirroring the decrease of the signal of micelles of CTAB in solution.

A more detailed analysis can be obtained through the study of the parameters extracted from the fits and given in Figure 3. This allows a sketch of the different steps of the reactions to be proposed, shown in Scheme 1.

The shape of micelles clearly evolves during the reaction, as seen from the values of the micelle radius R and the micelle ellipticity ε with time (Fig 3a and b). At $t=0$, micelles of CTAB can be fitted using a model of slightly prolate ellipsoids ($R=2.4\pm0.1$ nm and $\varepsilon=1.8\pm0.1$) in solution (step 1 in Scheme 1). With time, changes in the SANS pattern can be reproduced well by an elongation of the micelles ($R=2.2\pm0.1$ nm and $\varepsilon=3.0\pm0.1$ at $t=t_{\text{Bragg}}$), (step 2 in Scheme 1). After t_{Bragg} , the contribution of the micelles in the SANS pattern is modelled only by decreasing $I_{\text{micelles}}(q=0)$, without further modification of the micelles form factor (R and ε fixed). This elongation can be explained by the incorporation of silica species in the micelles. Indeed, from $I_{\text{micelles}}(q=0)$, we can extract the number of micelles per unit volume n and the scattering length density of the micelles ρ_{micelles} (Fig 3 c and d). Before precipitation ($t < t_{\text{Bragg}}$), we make the hypothesis that the number of micelles remains roughly constant in solution, assuming the change in shape can be mostly related to the addition of silica in their corona (although it is possible that reshaping of the micelles would induce a change in aggregation number and probably a slight decrease of n , neglected here). We can hence extract ρ_{micelles} , that is found to increase from $-0.242 \times 10^{-6} \text{ \AA}^{-2}$ at time $t=0$ (scattering length density of CTAB) to $0.9 \times 10^{-6} \text{ \AA}^{-2}$ once the Bragg peak is detected. From ρ_{micelles} , it is also possible to derive the amount of silica species x_{silica} in the micelles (Fig 3e), assuming micelles are composed only of CTAB and silica. This amount is found to reach almost 30% before the mesophase precipitation. As only a core model is needed to fit the data, it is impossible to describe with accuracy the localisation of the silica species in the micelles. Moreover, it is possible that a

fraction of D₂O is present in the micelles, neglected here, leading to a probable overestimation of x_{silica} . After t_{Bragg} , as the shape of the micelles does not evolve, $\rho_{micelles}$ is fixed while the decrease of $I_{micelles}(q=0)$ can be well described by the decrease of the number of micelles in solution n , consistent with their consumption in the particles to form the 2D-hexagonal network. The fact that this parameter equals zero at long time is a proof that most of the micelles are eventually consumed to form the mesophase (steps 3 and 4 in Scheme 1).

As seen on Figure 2, at $t=t_{Bragg}$ a signal of μ m-sized particles is detected in SEMSANS, concomitant with a Bragg peak of a mesophase in SANS.

The radius of the large particles $R_{particles}$ quickly reaches a very large size (at $t=t_{Bragg}=110$ min, fits give a $R_{particles}$ at ca 80 nm, while it is found above 2 μ m for the next measurement made at 129 min). Then, it slightly decreases upon time, again with some fluctuations, reaching 1.5 ± 0.2 μ m at 386 min (Fig 3f). As for the 2D-hexagonal organisation, the lattice parameter a does not vary significantly with time, whereas the apparent decrease in the domain size D observed around t_{Bragg} is probably explained by an overestimation of D at those early stages, due to the weak Bragg peak intensity at this time (Fig 3g and h). A value of D of approximately 80 nm is found throughout the entire kinetics series, which is more than ten times smaller than the particle radius observed in SEMSANS. This measurement of domain size may be artificially limited by the resolution of the SANS detector, however earlier measurement on film growth using X-ray reflectivity, where peak width was not defined by the detector resolution, suggested a domain size around 200 nm.³⁹ This size is still much smaller than the particle sizes observed here, which may suggest that the 1.5 μ m particles are composed of several 2D-hexagonal domains.

As described previously, after $t=t_{Bragg}$, we observed three phenomena: the appearance of particles in the SEMSANS signal, the appearance and slow growth of the Bragg peak

associated with the 2D-hexagonal mesophase and the slow decrease of the signal from micelles in solution. The concomitance between these phenomena is a direct indication that the nucleation of the particles is due to the rapid aggregation of silica-enriched micelles into phase separated particles. This immediately triggers a 2D-hexagonal organisation of the micelles within the particles, due to the higher concentration and interactions between micelles. Interestingly, while Σ_t quickly reaches a plateau around $ca\ t=160$ min, indicating that most phase-separated particles are formed at that time (step 3 in Scheme 1), both $I_{micelles}(q=0)$ and I_{10} continue to evolve with time; which suggests that the micelle organisation continues for a long period after particle formation, by the formation of new 2D-hexagonal domains in the particles, as the domain size does not increase with time (step 4 in Scheme 1). Around 160 min where most particles are formed, the number of micelles which are not organised is ca half the initial number of micelles. These unorganised micelles could be either already present in the particles (hence at a relatively high concentration) or still free in solution (in a dilute regime). As the signal of unorganised micelles is always fitted as non-interacting, it is assumed that most are still in solution. This other half of unorganised micelles is consumed over time, mirroring the growth of the Bragg peak intensity I_{10} , suggesting that the increase of number of 2D-hexagonal domains in the particles is probably not only due to micelles reorganisation within the particles but also to further adsorption of more silica-enriched micelles from the water phase. Adsorption of surfactant-silica composite micelles onto silica surfaces in an epitaxial fashion is well-established^{11, 42} and once the particle surface exists it is highly likely that further accretion of micelles onto this surface will continually occur. Possibly, the decrease in size of the large particles arises due to the continuing silica condensation and re-organisation of both silica and micelles in the particles, leading to compaction of initial aggregates which are composed of several different ~ 80 nm domains of 2D hexagonal organisation.

Finally, after $t=160$ min, the polydispersity in size of the large particles is quite narrow, especially considering the fact that they are probably composed of several domains as already seen by the difference between the domain size and the particles radius. This reflects the formation process which is reminiscent of a phase separation, similar to coacervation observed in polymer-surfactant systems where uniform liquid droplets form at a critical composition.⁴³ A coacervation-like mechanism has previously been suggested for the formation of micelle templated silica particles formed in bulk solution in systems where no film formation occurs.⁴⁴

The mesostructured silica film at this silica:surfactant ratio is hence obtained by later migration of a portion of the 2D-hexagonal ordered particles to the interface (step 5 in Scheme 1).

Increasing the amount of TMOS (CTAB, $r_{mol}=7.21$ and 10.81)

Two other syntheses using CTAB as surfactant were carried at higher TMOS concentrations ($r_{mol}=7.21$ and 10.81 , in Figures 4 and 5 respectively, and also in S.I. S3, S4 and S5). Similar stages to the kinetics were also observed when $r_{mol}=7.21$ compared to the already discussed $r_{mol}=3.6$ (SANS and SEMSANS data are given in Figure 4). Nonetheless, observation of μ m-scaled particles is observed at an earlier time for $r_{mol}=7.21$ (already at 66 min, compared to 110 min for $r_{mol}=3.6$), see Figure 6 and Figure S6, S.I. A smaller particle radius is also found, at approximately $1.1 \mu\text{m}$, whereas the lattice parameter and domain size are similar to the case $r_{mol}=3.6$ (see Figure S7, S.I.). The fact that precipitation occurs sooner at $r_{mol}=7.21$ is expected as more silica species are present.²⁰ Also, this fastest precipitation is associated with micelles less enriched with silica (x_{silica} around 15%, compared to 30%) and less anisotropic (at the precipitation time, $\varepsilon=2.5\pm0.1$ for $r_{mol}=7.21$, compared to 3.0 ± 0.1 for $r_{mol}=3.6$, see Figures 3 and 6). This suggests that the precipitation time is driven by the silica polymer size rather than

the micelle properties, similar to the effects of polymer molecular weight on coacervation observed in polymer surfactant systems. In such systems a threshold molecular weight must be achieved before phase separation into droplets occurs,⁴³ and here, as silica concentration increases this molecular weight will be reached more rapidly.

For the molar ratio $r_{mol}=10.81$, a loss of the beam prevented us to measure the beginning of the kinetic run (see SANS and SEMSANS measurements in Figure S4 and S5, S.I.). Although a signal from SEMSANS shows the presence of ca 1-2 μm particles once the beam was restored (first pattern at 164 min, see Figure S4), no Bragg peak can be seen in the highly noisy SANS patterns, in correspondence with earlier SANS studies on solutions at this ratio.³⁶ Hence, to have a better understanding of the mechanisms involved in that reaction, we performed additional SESANS experiments at Delft, and compared them to a set of data previously measured using SANS on LOQ.³⁶ SESANS and SANS data are given in Figure 5. The most striking aspect is that in this case, even if CTAB micelles are very slowly elongating upon time (see fitting parameters given in Figure 7), no Bragg peak is observed. Moreover, at long times (around 650 min in SANS and 800 min in SESANS), a sudden change in the signal is observed in SANS and a signal of very large particles (ca 8 μm) is detected in SESANS. The modification of the SANS signal gives a sharp increase of both the scaling factor and the ellipticity of the micelles (see Figure 7), but it is believed this is an artefact due to the growth of those large objects measured in SESANS. Indeed, the fits obtained at those times cannot reproduce accurately the data, with the emergence of a “shoulder” around 0.1 \AA^{-1} . This shoulder, associated with the increase of the signal at small angle ($q < 0.05 \text{ \AA}^{-1}$), could be the signature of the formation of disordered fractal-like silica gel particles, possibly from further aggregation of the 1-2 μm particles observed in SEMSANS at earlier times. We note that below 300 min, the CTAB micelles in this system are still only slightly prolate, with an ellipticity < 1.5 (see Figure 7). This would indicate that if the phase-separation is triggered by

the silica polymer size, the rearrangement into a mesophase is controlled by the micelle properties and most importantly its silica loading. Associated with the absence of Bragg peaks, these results indicate that for this ratio, a silica polymer network is forming in solution almost independently of the CTAB micelles, associated with the high amount of silica species in the solution, favouring silica-silica over silica-CTAB interactions. This silica network does collapse into particles rather than forming a solution spanning gel, trapping the micelles but at this point the silica polymer is too condensed to allow rearrangement into the ordered 2D hexagonal phase within the particles. The difference in times between the changes observed in SANS (650 min) and SESANS (800 min) is not necessarily significant as small variations in reproducing the same synthesis (*e.g.* slight changes in pH or level of deuteration) might influence the times associated with each event in the kinetics.

Those results are a strong indication that the mechanisms of the film formation are profoundly modified at this ratio, which is correlated with the fact that films formed much more slowly at $r_{mol}=10.80$ with CTAB have a much smoother interface compared to $r_{mol}=3.6$ and 7.21 .²⁰ The difference of roughness of the film has been previously explained by suggesting that for $r_{mol}=10.81$, it was formed by the aggregation of individual micelles at the interface, whereas for the other ratios the film is formed by already formed 2D-hexagonal particles migrating to the interface. This scenario is consistent with the results obtained in this study, especially the absence of mesostructured particles at $r_{mol}=10.81$. It is possible that the high amount of silica species creates a competition between adsorption of silica species onto the micelles and silica-silica condensation, preventing the organisation of the elongated micelles into a 2D-hexagonal order within particles in the bulk. The elongated micelles will hence more slowly migrate to the interface, to form a smooth silica film, with a 2D-hexagonal organisation to allow the highest packing between the micelles. On the contrary, for the two other ratios, the rapid formation of mesostructured silica particles in the subphase consumed the hybrid micelles, as

seen by the decrease of the number of micelles per unit volume n , and the film is formed by accumulation of these submicrometer grains at the interface. The small amount of mesostructured grains involved in the film formation compared to the ones present in the subphase prevents the quantification of this phenomenon.

Comparison with a synthesis made using $C_{16}EO_8$ at $r_{mol}=7.21$

A final synthesis was studied replacing CTAB by $C_{16}EO_8$, at a molar ratio $r_{mol}=7.21$ (Figure 8). Before addition of TMOS, $C_{16}EO_8$ micelles can be well modelled by a purely spherical model with a radius $R=3.2\pm0.1$ nm. However, similarly to the synthesis with CTAB, an elongation of the micelles is observed in solution upon time (until $t=110$ min), with a micelle radius decreasing from 3.2 to 2.5 nm, while the ellipticity ε increase from 1 to 4.8 (see parameters in Figure 9). $C_{16}EO_8$ micelles experience more drastic changes in their morphology upon the synthesis than CTAB at the same ratio, which is expected due to the larger headgroup that will interact with the silica species (8 EO groups compared to $N(CH_3)_3^+$). This results in a higher proportion of silica within the micelles (above 50% compared to 15% for CTAB at the same ratio) and an overall slower kinetics compared to CTAB at this silica:surfactant ratio.

At time $t=110$ min, a Bragg peak arises associated with the precipitation of the mesostructure in the subphase, once again concomitant with the appearance of a signal from particles in SEMSANS (see parameters in Figure 9). The same events as previously observed with CTAB can be pinpointed with this system, even if 2 μm sized particles and ordered mesophase are still forming at the last recorded pattern (239 min), evidencing a slower evolution than for CTAB at the same ratio.

As only one Bragg peak is observed, questions about the type of mesostructure formed in the particles arise. The peak is found at position $q=0.110\pm0.005$ \AA^{-1} which gives a d-spacing

$d=5.7\pm0.2$ nm. Previous studies of films made in these synthesis conditions showed a $Pm\bar{3}n$ cubic mesophase in the films, but with a d-spacing expected at 6.3 nm²¹. A d-spacing of 5.7 nm was found for C₁₆EO₈ mesostructured films at a ratio $r_{mol}=3.6$ and associated with a 2D-hexagonal mesophase.

The results demonstrate that the formation of micelle templated silica particles CTAB and C₁₆EO₈ follow the same steps, characterised by an elongation of the micelles due to the incorporation of silica species, before phase separation and formation of the mesostructure, observed simultaneously in SANS and SEMSANS. It is expected that for $r_{mol}=3.6$ and 7.21, these particles slowly migrate to the interface to form the film, as previously suggested.^{20-21, 39}

Conclusion

Using a combination of SEMSANS and SANS, kinetics of formation of mesoporous films were studied. Specifically the method allows measurement of the evolution of the system within the subphase over a wide range of length scales, in contrast to GISAXS or reflectivity that focus on the air-water interface. It was observed that for these syntheses, micelles of surfactant are initially present within the subphase. Evolution of the system is triggered by the addition of the silica source; which gradually hydrolyses in the acidic environment. This allows the incorporation of silica species within the micelles. Over time, incorporation of the silica species gradually elongates the micelles shape and the silica polymer grows in the solution. Finally, phase separation occurs, and mesostructured particles are formed in solution. Particles are formed containing micelles in an ordered mesophase – presumed to be a 2D hexagonal arrangement when CTAB is used as the templating surfactant – as can be observed by the concomitance of the emergence of a signal of particles in SEMSANS with the Bragg peak associated with the micelle organisation in SANS. The fact that no large objects can be observed in SEMSANS prior to the emergence of the Bragg peak in SANS, coupled

with the fact that micelles are non-interacting prior to the phase separation, suggests that upon precipitation micelles in the particles quickly rearrange in a mesophase with several domains but that the number of ordered domains increases with time due to further adsorption of silica enriched micelles, while the particle size shrinks a little due to the rearrangement of the micelles and continuing silica condensation. Indeed, after the formation of the particles, the signal of the remaining micelles in solution continues to decrease over time as they are added to the already formed particles. A portion of these mesostructured particles are assumed to migrate to the interface to form the rough mesostructured silica film previously observed in GISAXS. This model of formation of mesostructured particles is valid for both CTAB and C₁₆EO₈. The ratio of silica source compared to surfactant influences the reaction rate, or even disturbs this mechanism when too much silica is added, by competition between silica-surfactant and silica-silica interactions.

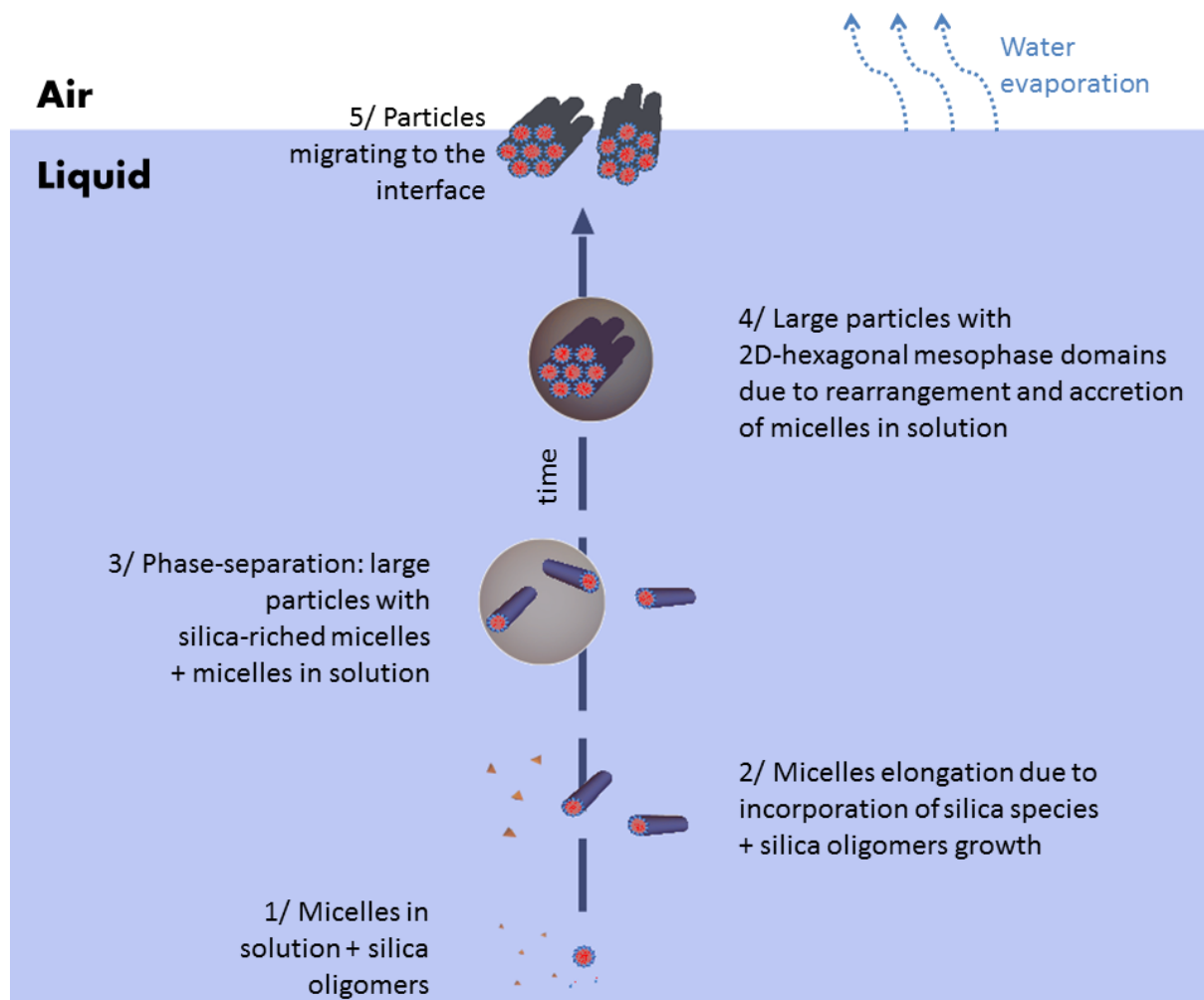
Acknowledgements

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Figures



Scheme 1: Sketch of the different steps proposed to be involved in the formation of a mesostructured film in the reaction using CTAB as surfactant at a silica:surfactant molar ratio

$$r_{mol}=3.6$$

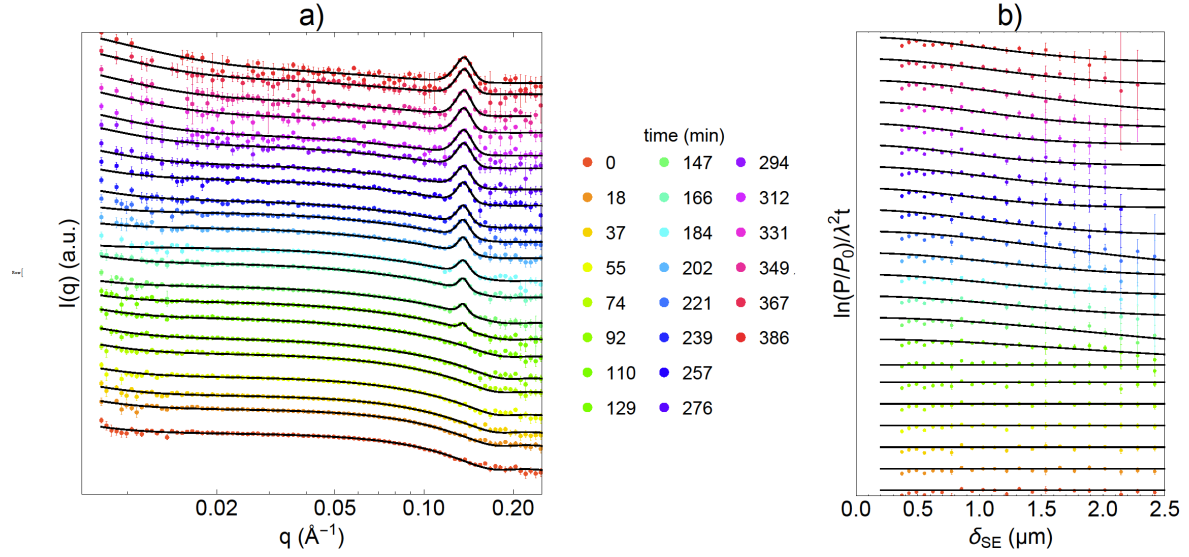


Figure 1: (a) $I(q)$ versus q (in \AA^{-1}) obtained from SANS measurements and (b) Normalized neutron spin-echo signal $\frac{\ln(P/P_0)}{\lambda^2 t}$ versus the spin-echo length δ_{SE} (in μm) obtained from SEMSANS measurements for the kinetics using CTAB as surfactant with $r_{mol}=3.6$. All patterns (a and b) have been shifted for clarity (non-shifted patterns can be found in Figure S1, S.I.) The colour gives the time associated with each pattern. In black lines, fits of the SANS and SEMSANS data are presented according to the models given in the Materials and Methods section.

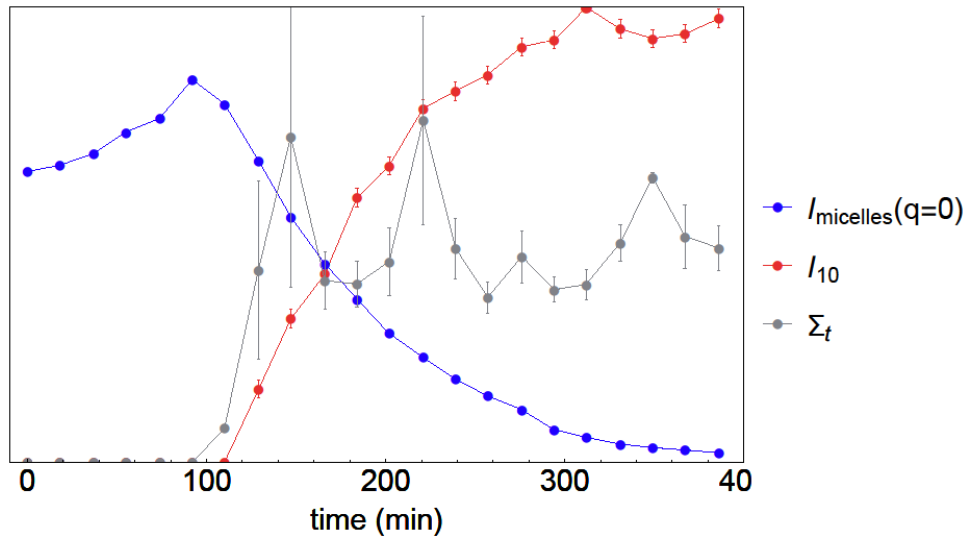


Figure 2: Comparison over time of the different contributions to the SANS and SEMSANS signals during the reaction in solution using CTAB at a ratio $r_{mol}=3.6$: in blue, the intensity $I_{micelles}(q=0)$ of CTAB micelles in solution extracted from the fitting of the SANS data, in grey the scattering power Σ_t of μm -sized particles measured in SEMSANS and in red the intensity I_{10} of the Bragg peak observed in SANS. Different scales in y have been overlapped to allow a clear depiction of the variation of these parameters with time. Figure S2 in S.I. gives separate graphs of these three parameters with their associated scales.

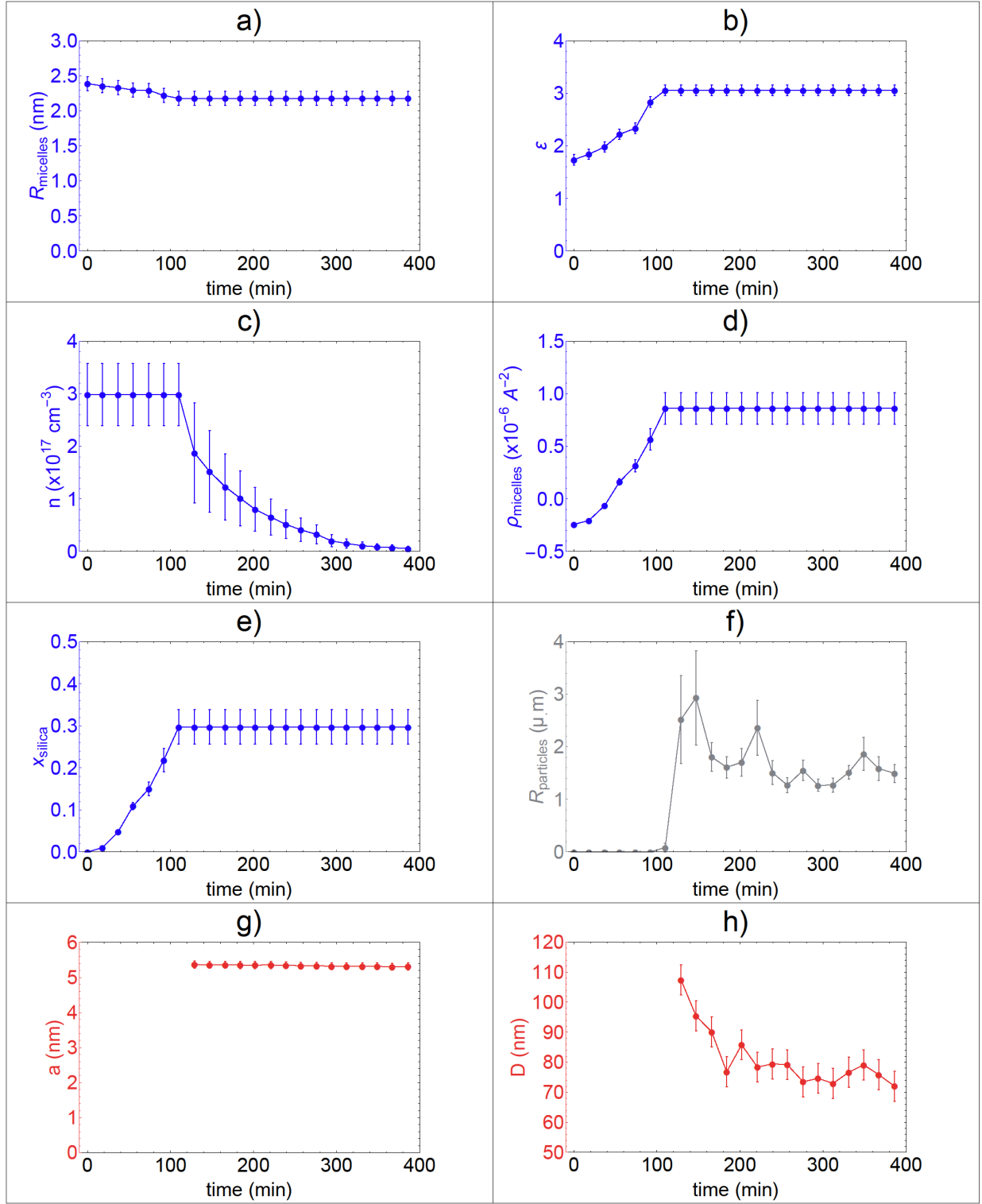


Figure 3: Evolution of different parameters extracted from fitting the SANS and SEMSANS data from the reaction in solution using CTAB at a ratio $r_{mol}=3.6$. In blue, parameters associated to the micelles of CTAB in solution: (a) the micelle radius $R_{micelles}$ (nm), (b) the

ellipticity of the micelles ε . From $I_{micelles}(q=0)$ (cm^{-1}) plotted in Figure 2, (c) the number of micelles per unit volume n (cm^{-3}) and (d) the scattering length density of the micelles $\rho_{micelles}$ (\AA^{-2}) were extracted, assuming that before the Bragg peaks appearance, n stays constant, while once the mesophase is forming, it is $\rho_{micelles}$ which does not change. (e) The proportion of silica within the micelles x_{silica} is extracted by $\rho_{micelles} = x_{silica}\rho_{silica} + (1 - x_{silica})\rho_{CTAB}$. In grey, (f) the radius R_{part} (μm) of μm -sized particles detected in SEMSANS. Finally in red, parameters associated with the 2D-hexagonal organisation within the μm -sized particles from fitting of the SANS Bragg peak: (g) the lattice parameter of the 2D-hexagonal order a (nm) and (h) the domain size D (nm).

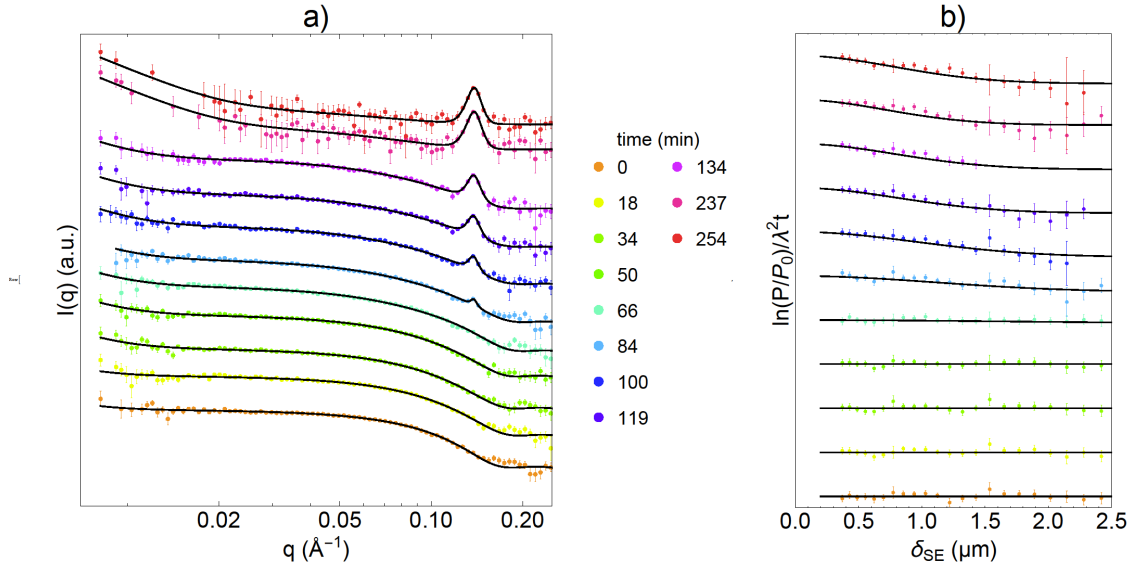


Figure 4: (a) $I(q)$ versus q (in \AA^{-1}) obtained from SANS measurements and (b) Normalized neutron spin-echo signal $\frac{\ln(P/P_0)}{\lambda^2 t}$ versus the spin-echo length δ_{SE} (in μm) obtained from SEMSANS measurements for the kinetic runs using CTAB as surfactant with $r_{mol}=7.21$. All patterns (a and b) have been shifted for clarity (non-shifted patterns can be found in Figure S3, S.I.) The colour gives the time associated with each pattern. In black lines, fits of the SANS and SEMSANS data are presented according to the models given in the Materials and Methods section.

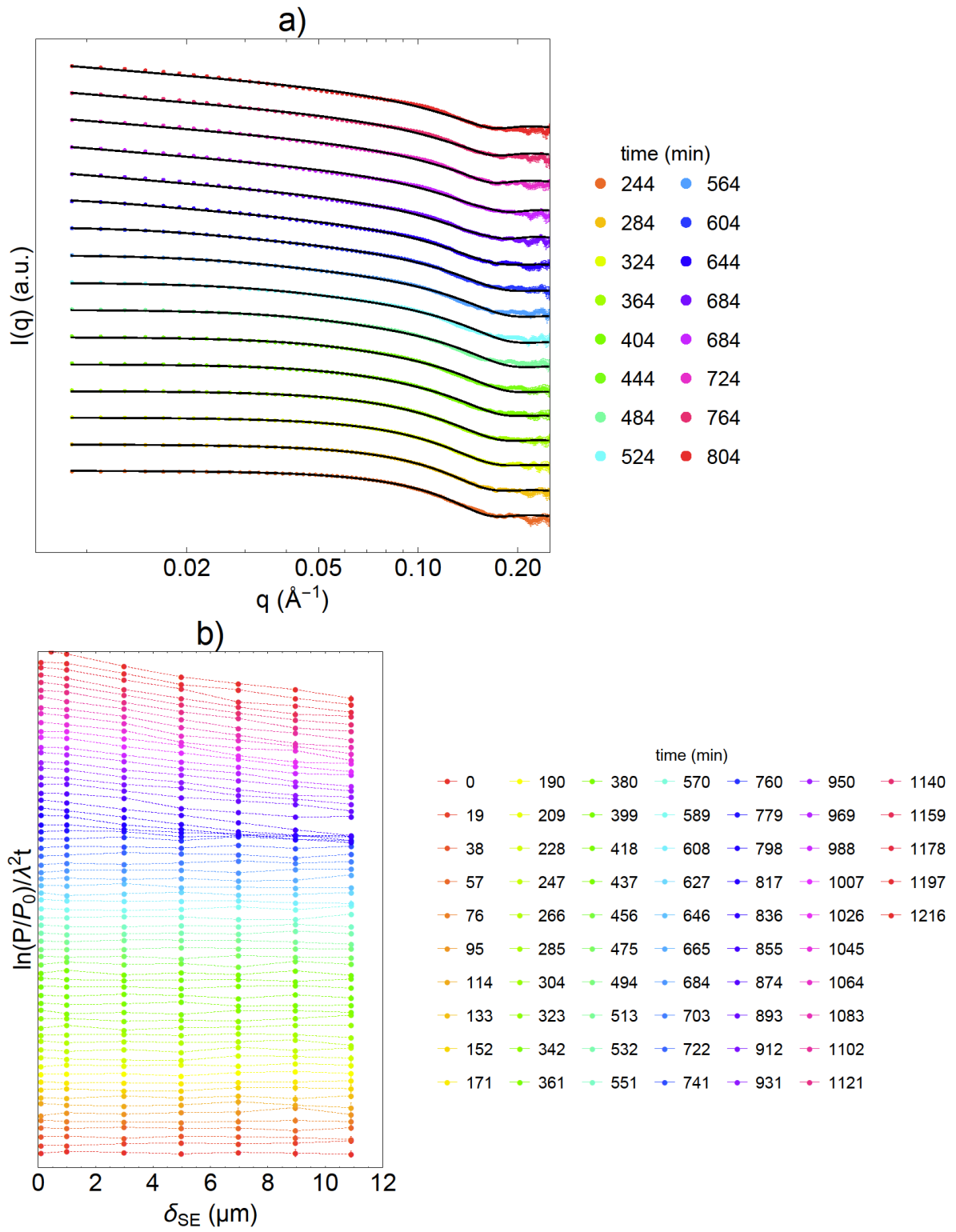


Figure 5: (a) $I(q)$ versus q (in \AA^{-1}) obtained from previous SANS measurements (on LOQ)³⁶ and (b) Normalized neutron spin-echo polarization $\frac{\ln(P/P_0)}{\lambda^2 t}$ versus the spin-echo length δ_{SE} (in μm) obtained from SESANS measurements at Delft for the kinetic runs using CTAB as surfactant with $r_{mol}=10.80$. Fits of the SANS data are provided in black lines.

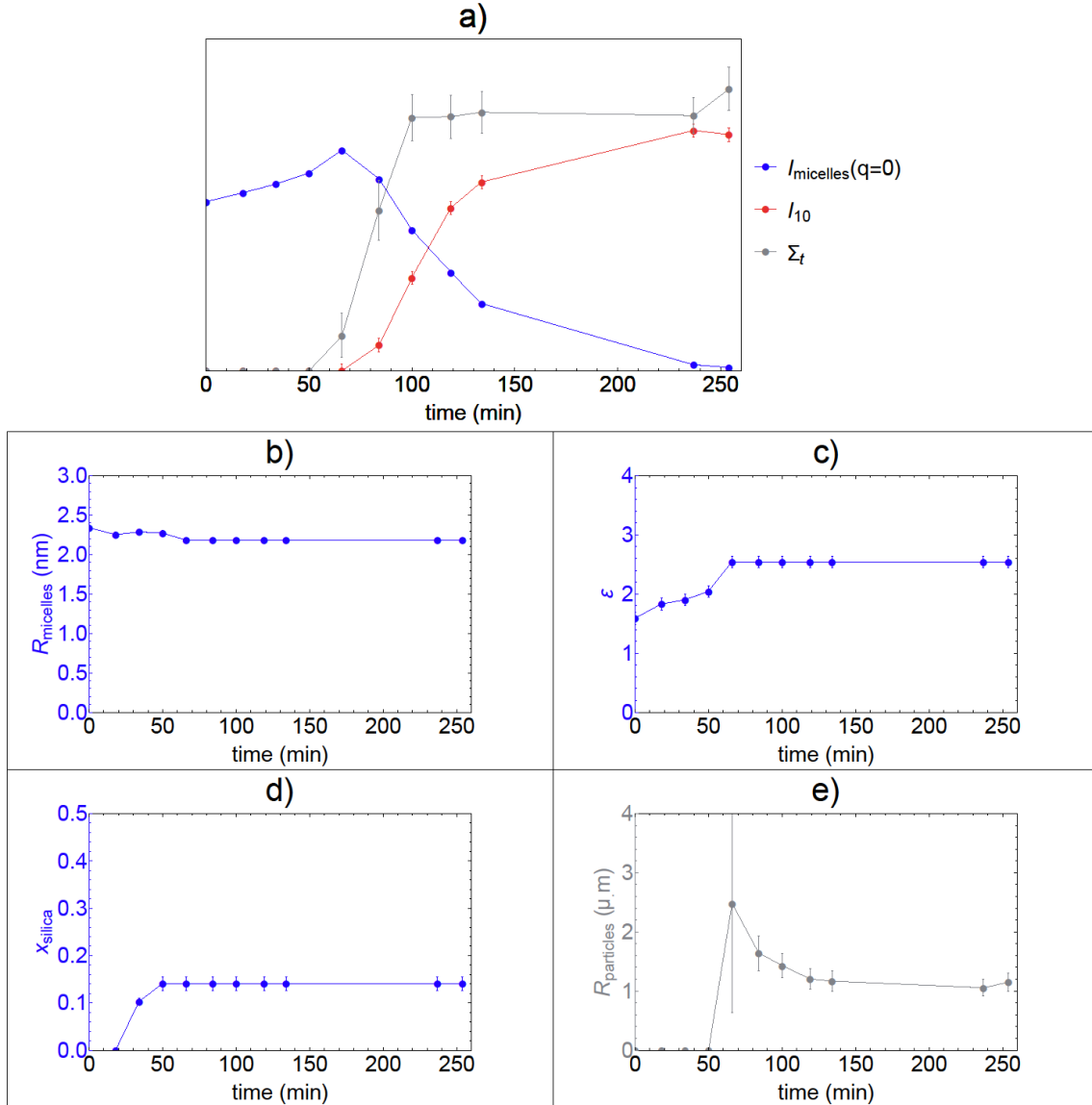


Figure 6: Evolution of different parameters extracted from fitting the SANS and SEMSANS data of the reaction in solution using CTAB at a ratio $r_{mol}=7.21$. (a) Comparison between $I_{micelles}(q=0)$ (in blue), I_{10} (in red) and Σ_t (in grey) upon time. Separated graphs with their associated scales can be found in Figure S6, S.I. In blue, (b) the micelle radius $R_{micelles}$ (nm),

(c) the ellipticity of the micelles ε and (d) x_{silica} (%) the amount of silica incorporated in the micelles obtained from fitting the SANS data. In grey, (e) the radius R_{part} (μm) of μm -sized particles detected in SEMSANS. Other fitting parameters can be found in Figure S7, S.I.

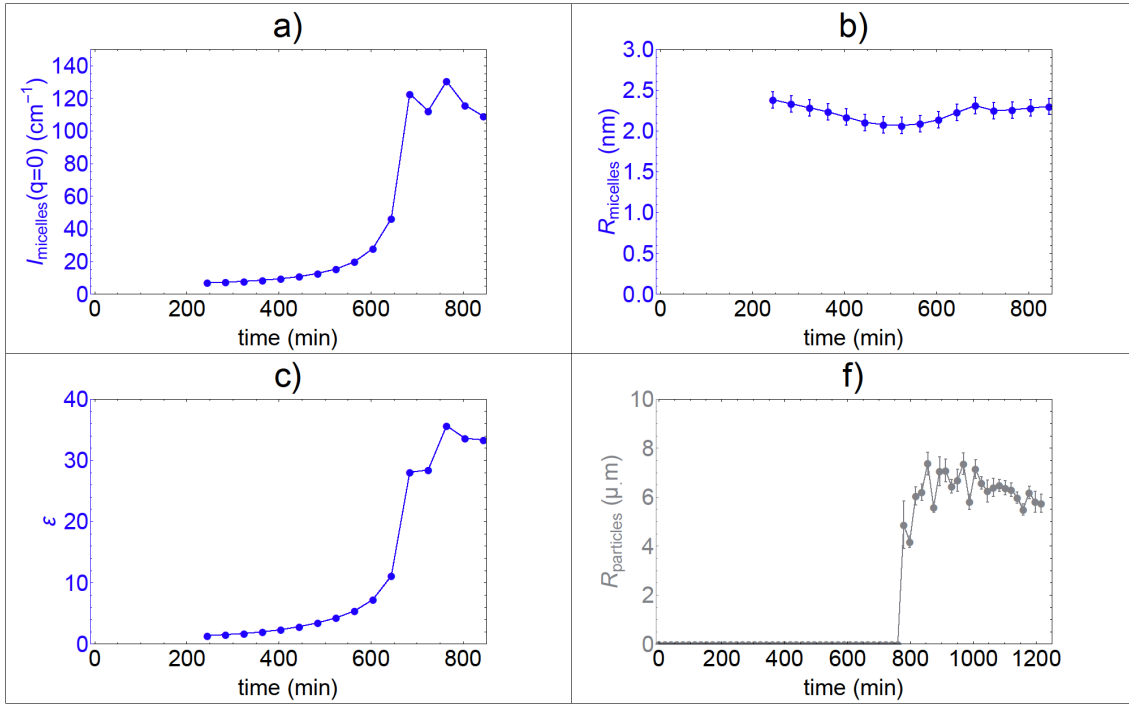


Figure 7: (a to c) Evolution with time of (a) $I_{\text{micelles}}(q=0)$ (cm^{-1}), (b) the micelle radius R_{micelles} (nm) and (c) the ellipticity of the micelles ε extracted from the fitting of the SANS data measured previously at LOQ. Around 650 min, a sudden increase in scale and ellipticity is observed, associated with the formation of a gel. (d) Particle radius measured in SESANS (Delft) over time. At 800 min, a signal of very large particles is observed and corresponds to macroscopic phase separation.

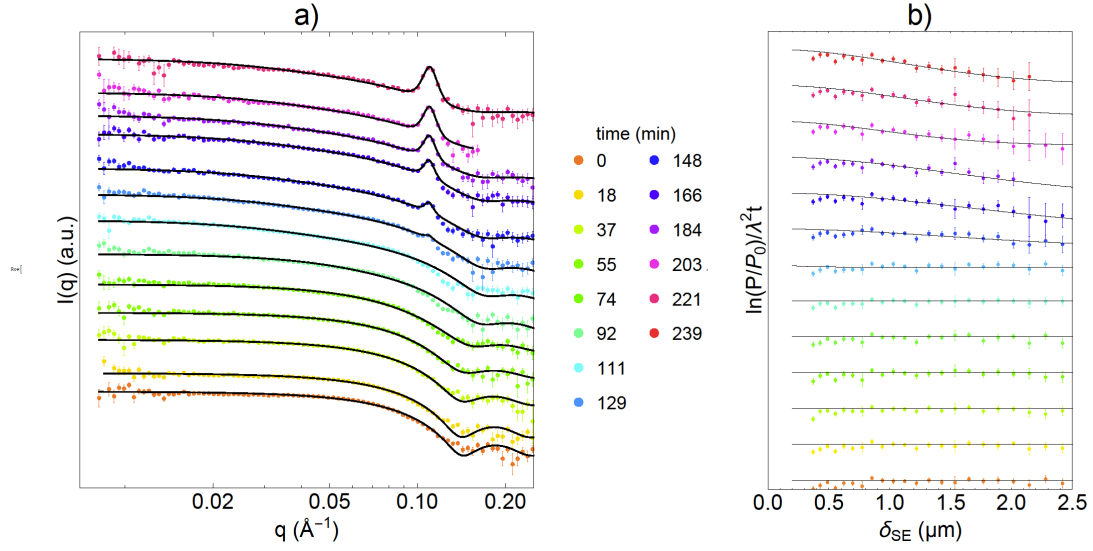


Figure 8: (a) $I(q)$ versus q (in \AA^{-1}) obtained from SANS measurements and (b) Normalized neutron spin-echo signal $\frac{\ln(P/P_0)}{\lambda^2 t}$ versus the spin-echo length δ_{SE} (in μm) obtained from SEMSANS measurements for the kinetic runs using C₁₆EO₈ as surfactant with $r_{mol}=7.21$. All patterns (a and b) were shifted for clarity (non-shifted SANS spectra can be found in Figure S8, S.I.) The colour gives the time associated with each pattern. In black lines, fits of the SANS and SEMSANS data are presented according to the models given in the Materials and Methods section.

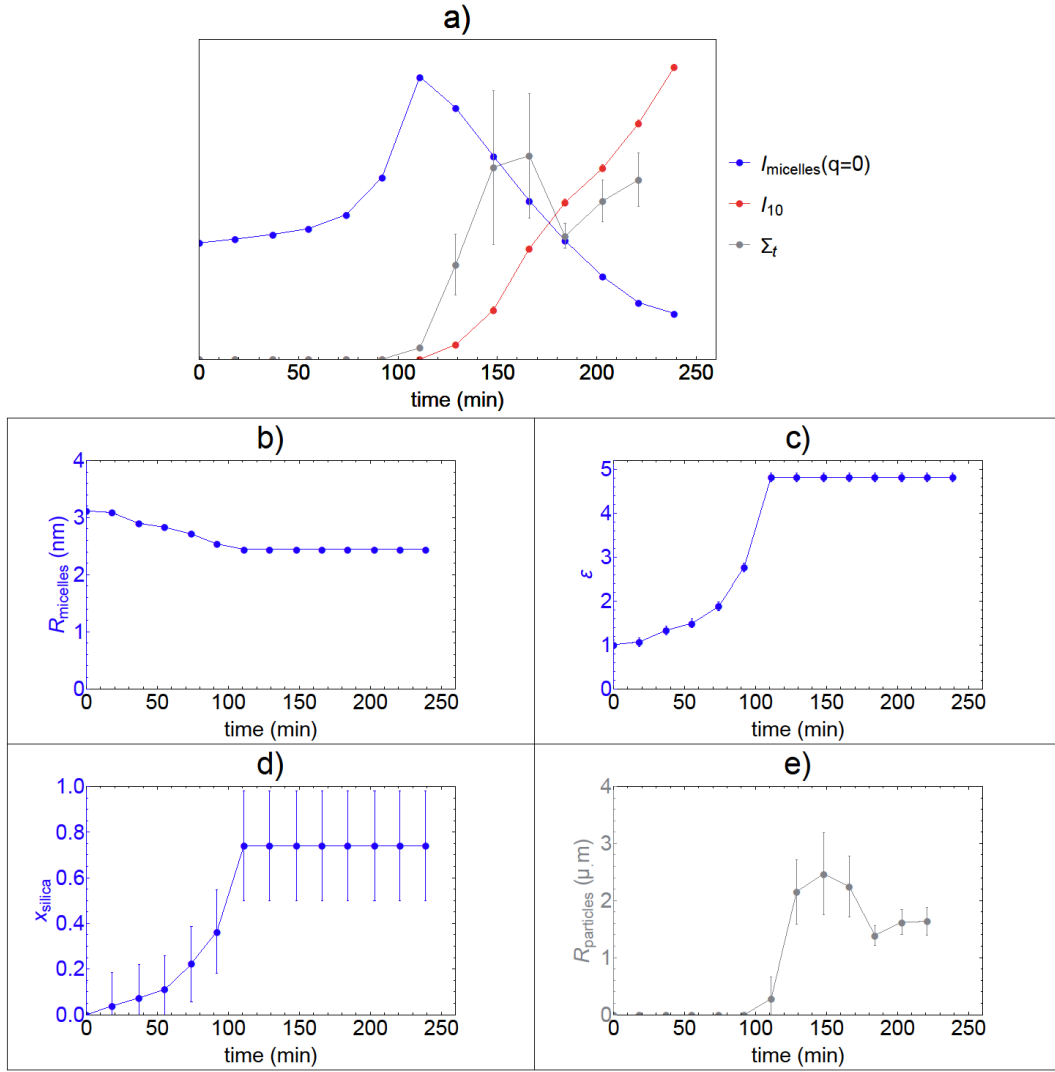


Figure 9: Evolution of different parameters extracted from fitting the SANS and SEMSANS data of the reaction in solution using $C_{16}EO_8$ at a ratio $r_{mol}=7.21$. (a) Comparison between $I_{micelles}(q=0)$ (in blue), I_{10} (in red) and Σ_t (in grey) upon time. Separated graphs with their associated scales can be found in Figure S9, S.I. In blue, (b) the micelle radius $R_{micelles}$ (nm), (c) the ellipticity of the micelles ε and (d) x_{silica} (%) the amount of silica incorporated in the micelles obtained from fitting the SANS data. In grey, (e) the radius R_{part} (μm) of μm -sized particles detected in SEMSANS. Other fitting parameters can be found in Figure S10, S.I.

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